

02/20/2001 10:04

(MON) 02/19/01 17:03 NO. 3561683386 P 2

FROM

## IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In Re the Application of:

HOWELL et al.

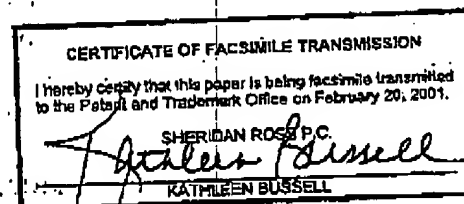
Serial No.: 09/444,144

Filed: November 20, 1999

Attorney File No.: 4369-1  
(formerly Cyto001)For: METHOD FOR ENHANCING  
IMMUNE RESPONSES IN  
MAMMALS

Group Art Unit: 1642

Examiner: Helms, L.

DECLARATION OF  
MARK D. HOWELL AND  
CHERYL L. SELINSKY  
(37 CFR § 1.132)RECEIVED  
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OFFICIAL

Assistant Commissioner for Patents  
Washington, D.C. 20231

Dear Sir:

We, Mark D. Howell and Cheryl L. Selinsky, each declare as follows:

1. I am a co-inventor of the above-referenced patent application and am familiar with the application.

2. This Declaration is being submitted in conjunction with an Amendment and Response After Final Rejection to an Office Action having a mailing date of December 20, 2000.

3. The following discussion is provided in traverse of the Examiner's rejection of Claims 1-3, 5, 10-27, 34, 37-38, 40-42, and 50-56 under 35 U.S.C. § 103. These comments have been discussed with the Examiner in the telephone interview of February 6, 2001, and this Declaration is provided at the Examiner's suggestion.

Discussion of Lentz

The Examiner contends that Lentz teaches the separation of blood into a plasma component and a cellular component. While Lentz acknowledges that it is possible to separate plasma from the blood, it is submitted that Lentz also *discourages* such a separation of blood (see col. 1, lines 51-62) on the basis that such a separation "has a serious impact on the platelet level in the blood," "could not be considered for widespread use," and is "not very attractive for clinical use." Therefore, the Lentz reference, at best, is highly ambiguous on this point.

02/20/2001 10:44

(MON) 02.19'01 17: ST. 17:03/NO. 3561683380 P 3

FROM

Second, a separation of cellular and acellular components offers no benefit if used in the method of Lentz. Lentz teaches a method of treating whole blood by passing the blood through a filter that separates factors in the blood based on size. Therefore, there is no advantage to be obtained by first separating the blood into cellular and acellular components, and then passing the components over the filter. Indeed, such a separation would be an added and unnecessary step in the method of Lentz. As such, the use of Lentz by the Examiner as a teaching of an element of plasma separation in the stated rejection under 35 U.S.C. § 103 is not appropriate because the teachings of Lentz are, at most, ambiguous, and further, because no advantage in the Lentz process is obtained by separating cellular and acellular components of whole blood. Therefore, Lentz does not provide any *motivation* to combine the ambiguous discussion in Lentz with the other cited references in the manner suggested by the Examiner.

In contrast to Lentz, and in further support of the non-obviousness of the method claimed in the present application, there are significant advantages not recognized or suggested by the prior art to separating the acellular fraction from the cellular fraction prior to treatment of the bodily fluid. First, it is noted that the present method has the advantage over Lentz of selectively removing the targeted immune system inhibitor without affecting the action of desirable immune system stimulators and other blood components. This advantage is significantly enhanced by treating only the acellular portion of the blood. Specifically, certain of the molecules targeted for removal by the method of the present invention are soluble components which typically bind to an immune system stimulator. These soluble components often are homologues of another binding partner for the immune system stimulator, such that the interaction between the immune system stimulator and the other binding partner is inhibited by the binding of the soluble component to the immune system stimulator. Since the other binding partner is frequently a cell-associated binding partner (i.e., is present on cell surfaces), it is desirable, and indeed, may be highly advantageous, to bind the soluble immune system inhibitor *without binding the homologous cell-associated binding partner*. For example, sTNFR1 is a soluble receptor for TNF $\alpha$  and TNF $\beta$ , which is produced through a proteolytic cleavage of the membrane receptor (mTNFR1) for TNF $\alpha$  and  $\beta$ . This proteolysis releases the extracellular domain of the mTNFR1 from the cell surface and allows it to diffuse freely into the extracellular space. The sTNFR1, thus produced, retains fully the ability to bind TNF  $\alpha$  and  $\beta$  with high affinity. Binding of TNF $\alpha$  and  $\beta$  by the sTNFR1 prevents TNF $\alpha$  and  $\beta$  from binding to the

FROM

mTNFRI. Consequently, the pro-inflammatory and apoptotic effects normally induced through the crosslinking of mTNFRI by TNF $\alpha$  and  $\beta$  also are inhibited. Due to the significant structural similarity between sTNFRI and mTNFRI, contact of whole blood with a binding partner reactive with sTNFRI would permit binding to both the sTNFRI in circulation and the mTNFRI present on cell surfaces. This would have dire consequences for the patient, and would contradict the goals of the present method for several reasons. First, binding of the binding partner to mTNFRI would block its engagement by TNF $\alpha$  and  $\beta$ , thus, effectively reducing TNF-induced immune responses. Second, binding of an immobilized binding partner to mTNFRI would effect the depletion of mTNFRI-bearing leukocytes from whole blood, thereby diminishing immune competence. Third, and most undesirable, binding of an immobilized binding partner to mTNFRI would crosslink the receptor and act, therefore, as an agonist of TNF $\alpha$  and  $\beta$ . This would produce very significant and potentially fatal toxicities similar to those observed in human clinical trials of infusional TNF $\alpha$ .

By separating the whole blood into acellular and cellular components in the claimed method, these issues are avoided and the advantages of selectively removing the targeted immune system inhibitor while maintaining the action of desirable immune stimulators and other blood components is achieved. Since Leutz does not teach or suggest any selective removal of any specific factor, these advantages can not be realized by the method of Leutz.

#### Discussion of Selinsky

The Examiner contends that the reference of Selinsky et al. teach that the soluble TNFRI is removed by Ultrapheresis [sic], and that with the knowledge of Leutz, one would know that soluble immune system inhibitors can be removed from whole blood. The Examiner has also pointed to the statement in Selinsky et al.: "[w]e, therefore, propose the development of methods and/or reagents capable of specifically removing sTNFRI, or antagonizing its effects *in situ*, as unconventional, yet promising, strategies for cancer immunotherapy."

It is submitted that, although the statement in Selinsky et al. may cause one of skill in the art to consider how to antagonize or remove sTNFRI *in situ*, such a statement is merely an invitation to experimentation and opens the door for one of skill in the art to consider a wide range of possible approaches. Indeed, Selinsky et al. provide absolutely no guidance as to how one of skill in the art would go about such a task, but rather generally state that the "therapeutic utility of manipulating

(MON) 02.19'01 17 P. 17:03/NO. 9561683386 P 5

FROM

sTNFRI levels *in vivo* has been demonstrated" and that "sTNFRI effectively inhibits immune responses *in vivo* and...its modulation is a legitimate therapeutic avenue." It is submitted that one of skill in the art, when presented with an invitation to manipulate the effects of a soluble protein, would look to a variety of conventional approaches to remove or manipulate the effects of that soluble protein *in vivo*, because such approaches are the most clinically desirable means of treating a patient. Conventional therapeutic manipulation of the immune system typically involves the administration of an antibody, peptide, protein, or small molecule that is designed to have a particular action in the patient. In fact, significant research has been directed to systems that enable the delivery of such reagents, including liposomes, targeting antibodies, combinations of liposomes and antibodies, small particles, emulsions, and other vehicles.

One conventional approach to modulating an immune response *in vivo* is to introduce into the subject a reagent that achieves the goal of selectively antagonizing or removing a target molecule once it is administered to a subject. For example, one method for removing or blocking the action of a soluble protein *in vivo* is to administer an antibody that binds to and effectively neutralizes the action of the target protein. Alternatively, a peptide or other soluble binding partner that competes with the target protein for binding to the natural ligand can be administered. As yet another alternative, a small molecule could be designed that targets and neutralizes the action of the target protein. Drug design for such *in vivo* applications is a common therapeutic approach when a target such as a soluble protein is available.

In contrast, to turn to an *ex vivo* approach such as that claimed in the present application is not conventional, and indeed, would be much less likely to be considered because it would conventionally be considered to be less direct, more expensive, and more invasive than the *in vivo* approaches discussed above. Such a method requires far greater manipulation of the patient and of the critical bodily fluids of the patient than an *in vivo* approach. Therefore, to arrive at the claimed *ex vivo* method would not have been an obvious extension of the statements made in Selinsky et al. that are referenced above.

4. I hereby declare that all statements made herein of my own are true and that all statements made on information and belief are believed to be true; and further that the statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both under Section 1001 of Title 18 of the United States Code, and that

FROM

(MON) 02/19/01 17:03/NO. 3501683380 P 6

such willful false statements may jeopardize the validity of the subject application or any patent  
issuing therefrom.

2/20/01  
Date

Mark D. Howell  
Mark D. Howell

2/20/01  
Date

Cheryl L. Selinsky  
Cheryl L. Selinsky